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LETTERS

Dynamics of DNA replication loops reveal temporal control of lagging-strand synthesis

Samir M. Hamdan¹, Joseph J. Loparo¹, Masateru Takahashi¹, Charles C. Richardson¹ & Antoine M. van Oijen¹

In all organisms, the protein machinery responsible for the replication of DNA, the replisome, is faced with a directionality problem. The antiparallel nature of duplex DNA permits the leading-strand polymerase to advance in a continuous fashion, but forces the lagging-strand polymerase to synthesize in the opposite direction. By extending RNA primers, the lagging-strand polymerase restarts at short intervals and produces Okazaki fragments^{1,2}. At least in prokaryotic systems, this directionality problem is solved by the formation of a loop in the lagging strand of the replication fork to reorient the lagging-strand DNA polymerase so that it advances in parallel with the leading-strand polymerase. The replication loop grows and shrinks during each cycle of Okazaki fragment synthesis³. Here we use single-molecule techniques to visualize, in real time, the formation and release of replication loops by individual replisomes of bacteriophage T7 supporting coordinated DNA replication. Analysis of the distributions of loop sizes and lag times between loops reveals that initiation of primer synthesis and the completion of an Okazaki fragment each serve as a trigger for loop release. The presence of two triggers may represent a fail-safe mechanism ensuring the timely reset of the replisome after the synthesis of every Okazaki fragment.

The 'trombone model' as proposed in ref. 3 provides an elegant model for coupling many rounds of Okazaki fragment synthesis on the lagging strand of the replication fork to the continuous production of DNA on the leading strand. The looping back of the lagging strand onto the replisome allows both leading- and lagging-strand DNA polymerases to synthesize in the same direction and facilitates recycling of the lagging-strand DNA polymerase by virtue of its proximity to the RNA primers newly synthesized at the fork. The visualization of replication intermediates of the T4 and T7 bacteriophage replication systems by electron microscopy demonstrated the existence of replication loops and allowed a characterization of their length distributions^{4,5}. Biochemical studies revealed a number of molecular scenarios that explain how formation and release of replication loops may be regulated^{6–14}. However, no dynamic characterization has been reported and the timeline controlling the various enzymatic activities at the fork is not entirely understood.

In this study, we report the reconstitution of T7 replisomes and the observation of coordinated leading- and lagging-strand synthesis at the single-molecule level. The T7 replisome consists of only four proteins and it displays all the important features of more complicated replication systems¹⁵ (Fig. 1a). These proteins are the T7 DNA polymerase, a 1:1 complex of T7 gene 5 protein (gp5) and *Escherichia coli* thioredoxin processivity factor, T7 gene 4 helicase–primase protein (gp4), and T7 gene 2.5 single-stranded DNA binding protein (gp2.5). In previous work, we used single-molecule techniques to study the activity of partially assembled replisome proteins mediating only leading-strand synthesis in both T7 and *E. coli*^{16–18}. Here we describe the observation of T7 replication complexes undergoing

coordinated leading- and lagging-strand synthesis, allowing for kinetic characterization of many rounds of replication loop formation and release. Replication is studied at the single-molecule level by monitoring the length of individual, tethered DNA molecules. Briefly, the lagging strand of a forked-duplex phage- λ DNA molecule (48.5 kilobases (kb) long) is attached by one end to the surface of a glass flow cell and the other end to a bead¹⁹ (Fig. 1b; Supplementary Information). A laminar flow exerts a well-controlled drag force on the beads and stretches the DNA molecules. Figure 1c shows the length of an individual DNA molecule as a function of time in the flow of a buffer containing purified gp4, T7 DNA polymerase, gp2.5, Mg^{2+} , four deoxynucleoside 5'-triphosphates and the ribonucleotides adenosine triphosphate (ATP) and cytidine triphosphate (CTP) required for primase activity¹³. The single-molecule trajectories show repeated cycles of DNA shortening (Fig. 1c, blue arrow) followed by rapid lengthening to the original DNA length (Fig. 1c, red arrow).

In the presence of a saturating concentration of gp2.5, which is required to coordinate leading- and lagging-strand synthesis^{13,20}, single-stranded DNA (ssDNA) is equal in length to double-stranded DNA (dsDNA) (Supplementary Information). Because the bead is attached to the surface by the lagging strand, the formation of a replication loop in this strand will decrease the apparent DNA length by an amount equal to the DNA length stored in the loop. Several lines of evidence support the notion that the observed DNA shortening is a result of coupled DNA replication and loop formation. First, the loop length is comparable to that observed in electron microscopy studies. Figure 1d shows the wide distribution of loop lengths observed for 115 individual replisomes. This distribution can be described using a single-exponential distribution with a decay constant of 1.4 ± 0.1 kb, which is close to the average length of 2 kb obtained in previous electron microscopy studies⁵.

The observation that inhibition of either the primase or DNA polymerase activity abrogates DNA shortening provides further confirmation that the observed events are related to coupled replication (Supplementary Information). Furthermore, the average rate of DNA shortening observed during loop growth (146 ± 50 bp s⁻¹; Fig. 1e) is nearly twice the rate observed for leading-strand polymerization alone under similar experimental conditions (80 bp s⁻¹) (Supplementary Information)¹⁶, consistent with the notion that loop growth is supported by both leading- and lagging-strand synthesis with a net rate that contains the contributions of the two polymerases. Also, we observe multiple DNA shortening events (an average of 3 ± 2 loops per replication event, with 25% of replisomes displaying more than 4 loops; Supplementary Information) that occur in rapid succession on a small fraction of surface-tethered DNA molecules (5%). This pattern indicates the presence of stably assembled and processive replisomes, as opposed to the distributive activity of replication proteins leading to isolated looping events. Also, loop length and the number of successive loops formed per

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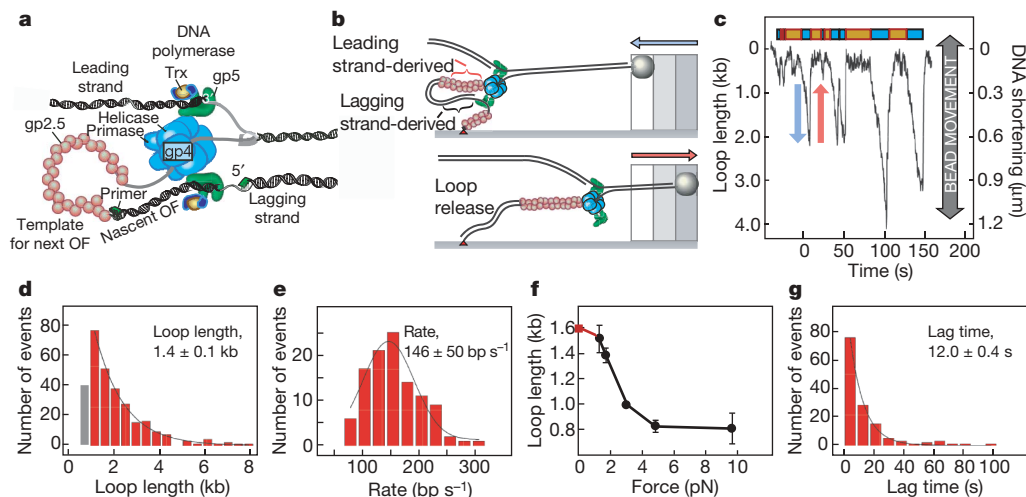


Figure 1 | Observation of replication loops. **a**, Organization of the T7 replication fork. Gene 4 protein (gp4) encircles the lagging strand and mediates both the unwinding of double-stranded DNA through its helicase domain and the synthesis of RNA primers through the primase domain. The T7 DNA polymerases are stably bound to gp4 and incorporate nucleotides on the leading and lagging strands. The DNA polymerase is a 1:1 complex of the T7 gene 5 protein (gp5) and *E. coli* thioredoxin (trx). The ssDNA extruded behind the helicase is coated by the ssDNA-binding protein gp2.5. A replication loop is formed in the lagging strand to allow the polymerase to synthesize in the same direction. The lagging-strand DNA polymerase initiates the replication of Okazaki fragments (OFs) using RNA primers synthesized by the primase domain of gp4. **b**, Depiction of the overall length change of tethered DNA as a result of replication loop formation and release. The blue and red arrows correspond to shortening and lengthening of the DNA; see **c**. **c**, Change over time of the length of a single DNA molecule during replication (see Supplementary Information for more examples). The loop growth and lag phases are shown as cyan and orange boxes, respectively.

replisome are reduced by increasing force (Fig. 1f). This observation is consistent with the prediction that the applied force will be exerted directly on the protein interactions that hold the loop together.

Finally, fluorescence time-lapse microscopy on individual, replicating DNA molecules demonstrates that DNA is synthesized on both the leading and the lagging strand (Fig. 2). In these experiments, λ DNA templates were stained with a fluorescent, dsDNA-specific dye and the fluorescent DNA was imaged while flow-stretched and replicated by the T7 replisome. The growth of a leading-strand product, its continuous movement along DNA, and the double-stranded nature of the lagging-strand product demonstrate that both leading and lagging strands are being replicated. Further confirmation of the presence of both leading- and lagging-strand synthesis is obtained by a bulk-phase analysis of replication products obtained under conditions identical to those used in the single-molecule experiments (Supplementary Information). The length of Okazaki fragments (0.8 kb) corresponds well with half of the mean loop length as measured in the single-molecule experiments ($0.5 \times 1.4 \text{ kb} = 0.7 \text{ kb}$), in agreement with the fact that half of the replication loop consists of the nascent Okazaki fragment produced by the lagging-strand DNA polymerase and the other half consists of ssDNA extruded by the helicase⁵.

Observation of replication loop formation and release provides us with kinetic information that is inaccessible using bulk-phase assays. The stochastic and sequential nature of the many enzymatic processes involved causes the synchronicity of a population of reactions to be lost quickly and will obscure kinetic information about the short-lived, intermediate steps. Our single-molecule experiments reveal the presence of lag times between the release of one replication loop and formation of the next, a step unobservable in bulk-phase assays. The distribution of the lag times follows a single-exponential dependence with a decay constant of $12.0 \pm 0.4 \text{ s}$ (Fig. 1g). The

respectively. **d**, Histogram of the replication loop length (bars; $n = 288$) fitted with a single-exponential decay (solid line). The grey bar represents loop lengths below 1 kb, which are under sampled owing to noise in the length measurements. These loops were not included in the fitting of loop size distributions. **e**, Histogram of rate of loop growth (bars; $n = 107$) fitted with a normal distribution (solid line). The rate of loop growth is determined from the slope of the DNA shortening phase. **f**, Dependence of replication loop length on stretching force ($n = 108, 288, 158, 64, 26$ at forces of 1.4, 1.7, 3.0, 4.8 and 9.5 pN, respectively). The replication loop length at zero force (red square) is derived from the Okazaki fragment length as measured in bulk-phase experiments (Supplementary Information). **g**, Histogram of lag times (bars; $n = 135$) fitted with a single-exponential decay (solid line). All reported mean values are obtained using the maximum-likelihood-estimation method. Uncertainties (**d**, **e**, **g**) and error bars (**f**) correspond to the standard deviation of the distribution (**e**) or to the error of fitting the loop length (**d**, **f**) and lag time (**g**) distributions with exponential decay functions.

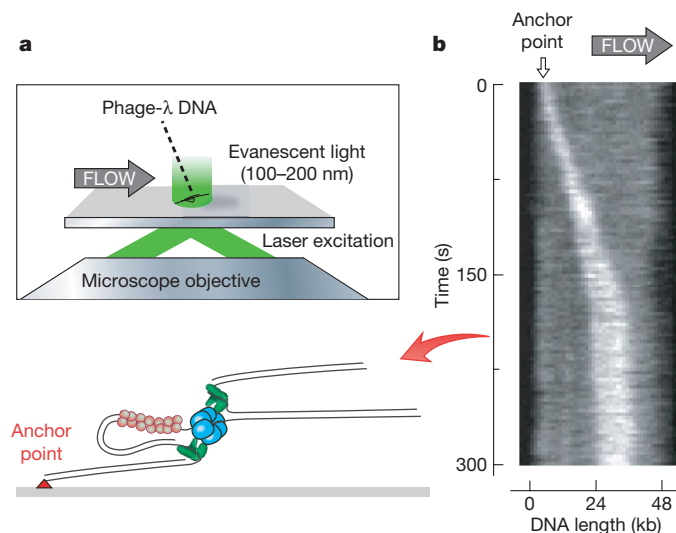


Figure 2 | Fluorescence imaging of coordinated DNA replication. **a**, The experimental design. Individual DNA molecules are fluorescence-stained by means of intercalating dye, stretched by flow and imaged through total-internal-reflection fluorescence (TIRF) microscopy. As the replication reaction proceeds, the leading-strand product will grow and be flow-stretched along the parental DNA. **b**, Kymograph of an individual DNA molecule undergoing coordinated replication (see Supplementary Movie). The grey scale indicates the fluorescence intensity. The intensity is doubled where the leading strand overlaps the parental strand. Growth of the leading strand and its movement along the DNA indicate processive replication. The double-stranded nature of the DNA between anchor point and leading strand indicate synthesis at the lagging strand.

appearance of a lag time after replication loop release suggests the requirement for intermediate steps before the formation of a new replication loop. At least three steps are necessary for a loop to be formed: the recognition by the primase of a priming site in the lagging strand, the synthesis of a primer and its handover to the lagging-strand DNA polymerase. The importance of primer synthesis in regulating the timing of the events at the replication fork is clear from several studies that found an effect of primase activity on the size of Okazaki fragments in T7, T4 and *E. coli* replisomes^{10,12,13}. However, the timing of primer synthesis and its causal relation with loop release has not been established.

By changing the concentration and nature of the ribonucleotides required by the primase, we can modulate the observed loop dynamics and extract a timeline of enzymatic events during the replication cycle. T7 DNA primase recognizes four sequences, namely 5'-GGGTC-3', 5'-TGGTC-3', 5'-GTGTC-3' and 5'-TTGTC-3', and requires only ATP and CTP to synthesize primers (5'-ACCC-3', 5'-ACCA-3', 5'-ACAC-3' and 5'-ACAA-3') complementary to the four nucleotides at the 5' end of the recognition sequence²¹. Reducing the ATP and CTP concentrations from the optimal 300 μ M to 30 and 10 μ M, respectively, shows an increase in loop length from 1.4 ± 0.1 kb to 3.6 ± 0.5 kb (Fig. 3). The lag time between loops also increases significantly, from 12.0 ± 0.4 s to 28.6 ± 2.8 s (Fig. 3). The observation that changing the kinetics of primer synthesis alters the loop lengths as well as the length of the lag times led us to consider how the different steps in primer synthesis have a role in the timing of the events that lead to both loop release and loop formation. Primer synthesis takes place in two distinct steps^{22–24}. First, the primase condenses ATP and CTP to form pppAC, the triphosphate diribonucleotide that is present as the starting nucleotide pair in all four possible primers. Subsequently, pppAC is extended in a much slower step to a full-length, tetranucleotide primer in a sequence-dependent manner^{22–24}. T7 DNA primase can utilize pAC, the monophosphate AC diribonucleotide, and extend it efficiently using only ATP and CTP²³. Therefore, by providing the coupled replication reaction with a pre-made pAC, we can bypass the requirement of the condensation step. At low ATP (30 μ M) and CTP (10 μ M) concentrations, the presence of 300 μ M pAC nearly restored the loop length to that

observed at optimal ATP and CTP concentrations (Fig. 3). The lag time, however, remains unaffected by the presence of pAC. These results demonstrate that the first step in primer synthesis, condensation of ATP and CTP to form pppAC, triggers loop release. As a consequence, the lag time has to include the slow extension step of pppAC to a full tetranucleotide.

Our observation that replication loop length is determined by primer synthesis and previously reported dependencies of Okazaki fragment length on primase activity^{10,12,13} suggests a signalling mechanism by which the primase controls the timing of the cycle of enzymatic events at the fork. The observation that the average Okazaki fragment length as observed in the bulk phase does not decrease on introduction of pAC (Supplementary Information) demonstrates that pAC triggers loop release before the nascent Okazaki fragment is completed, explaining previous observations of ssDNA gaps in electron microscopy and bulk-phase assays^{10,11}. Primer synthesis and loop release before completion of the Okazaki fragment will result in a length decrease of the ssDNA template available for the next Okazaki fragment. Consequently, a gradual decrease in replication loop length is predicted as the replisome progresses. However, a length comparison between subsequent replication loops formed by individual replisomes suggests no apparent trend in loop size (mean length difference is 0.17 ± 0.3 kb).

In an alternative model, the encounter of the lagging-strand DNA polymerase with the 5' terminus of the previously synthesized Okazaki fragment triggers dissociation of the polymerase and subsequent loop release^{6–9}. In this collision model, leading-strand synthesis needs to continue after loop release to allow the primase to search for its recognition sequence. In this case, the size of the subsequent Okazaki fragment is expected to increase by the extra amount of ssDNA generated during the primase search. Therefore, the collision model predicts a gradual increase in Okazaki fragment size as the replisome progresses. To address the possibility that both signalling and collision models are operative, preventing a net change in loop length, we test a number of predictions for the behaviour of the lag times between loops in the two different mechanisms. In both mechanisms, the lag time will contain the primer extension step and handover of the primer to the lagging-strand DNA polymerase. In the collision mechanism, however, the lag time will also include additional leading-strand synthesis mediating the search of the primase for a priming sequence. A lack of length contrast between ssDNA and dsDNA prevents us from directly observing leading-strand synthesis during the lag phase. Because further leading-strand synthesis will give rise to a lengthening of the Okazaki fragment produced during the next cycle and collision-mediated loop release requires completion of the fragment, we predict a positive correlation between the lag time and the increase in loop length. Similarly, we predict no correlation between lag time and loop length change in the case of a signalling mechanism. Inefficient primer utilization and a subsequent increase in single-stranded DNA template size may result in an increase in Okazaki fragment size, but loop release will still be triggered by the stochastic condensation of ATP and CTP and will remain uncorrelated with lag time.

To test the presence of both signalling and collision mechanisms, we divide all observed loop pairs into two categories, solely on the basis of whether the second loop of a pair is longer or shorter than the first loop of the pair, and determine the correlation between lag time and loop length change for each of the two groups. We see a correlation between lag time and loop length change in the data set that showed loop growth ($r = 0.55$, $n = 50$; the probability that a similarly sized data set of two uncorrelated variables would produce this correlation coefficient is less than 0.05%), but not in the group of loop pairs that showed a decrease in length ($r = -0.05$, $n = 49$) (Fig. 4). A slightly longer lag time between loop pairs that showed a length increase (collision controlled; 13.3 ± 1.5 s) and pairs that showed a length reduction (signalling controlled; 11.2 ± 0.8 s) lends further support to the notion that after collision-mediated release, extra leading-strand synthesis is required

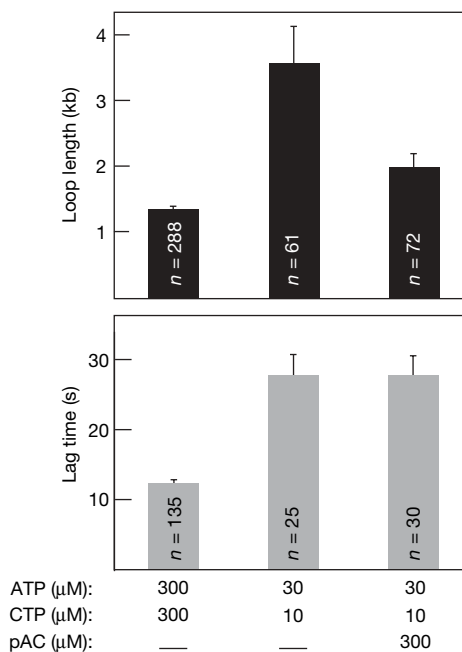


Figure 3 | Replication loop dynamics depend on primase activity. Loop growth and lag times are compared for different concentrations of ribonucleotides. Replication loop lengths and lag times were determined as described in Fig. 1.

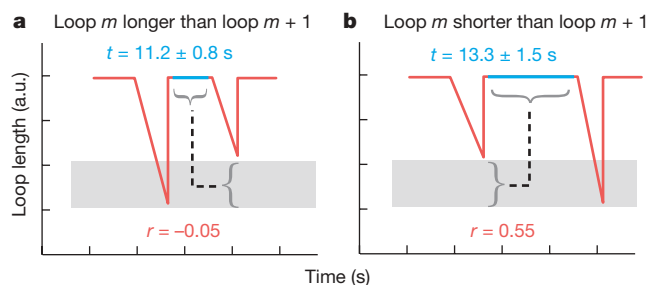


Figure 4 | Signalling and collision mechanisms both serve to trigger loop release. **a**, Correlation ($r = -0.05$, $n = 49$) between the lag time and the size change of the second loop for the pairs of loops in which loop $m + 1$ is smaller than loop m . **b**, Correlation ($r = 0.55$, $n = 50$) between the lag time and the size change of the second loop for the pairs of loops in which loop $m + 1$ is larger than loop m . a.u., arbitrary units.

to facilitate the search for a primer site. Additionally, experiments done under conditions that disfavour the collision mechanism by increasing the Okazaki fragment size while maintaining the pAC synthesis kinetics as a loop release signal result in the absence of any correlation between lag time and loop length (Supplementary Information). Taken together, these results suggest that the signalling and collision mechanisms are both operative during coupled replication.

In previous single-molecule studies, we have demonstrated that primase search is a stochastic process, with there being a limited probability of the primase recognizing and using a priming sequence during its scan of the lagging strand¹⁶. The random nature of primase activity poses a fundamental challenge to coordinating the timing of primer synthesis with the release of the replication loop. The utilization of both the signalling mechanism and the collision mechanism to release a replication loop and initiate formation of a new one provides an elegant mechanism that allows the replisome to cope with the stochastic nature of the primase activity. The signalling mechanism will release the replication loop if the primase locates one of its sequences before the nascent Okazaki fragment is finished. On the other hand, if the nascent Okazaki fragment is finished without the primase having engaged a recognition sequence, then the collision mechanism acts as a fail-safe mechanism to trigger loop release and ensure that the cycle of enzymatic events at the replication fork is properly reset.

METHODS SUMMARY

Single-molecule DNA stretching. A biotinylated fork was introduced at one end of phage- λ DNA to attach it to the streptavidin-coated glass surface of a flow cell. The other DNA end was functionalized with a digoxigenin moiety to couple it to a 2.8- μ m-diameter anti-digoxigenin-coated bead¹⁶. A laminar flow was introduced to exert a well-defined drag force on the bead, resulting in a stretching of the DNA molecules. Coordinated DNA synthesis was carried out in a flow of purified gp4 helicase–primase²⁵, T7 DNA polymerase²⁶, gp2.5²⁷ and nucleotides. Beads were imaged onto a charge-coupled device using dark-field microscopy and their positions tracked using particle-tracking software. For a detailed description, see Methods and Supplementary Information.

Fluorescence imaging. DNA was tethered at the forked end to functionalized coverslips as described above, but beads were omitted. In the presence of dsDNA-specific stain, the hydrodynamically stretched DNA was imaged using through-objective TIRF microscopy (Methods and Supplementary Information).

Full Methods and any associated references are available in the online version of the paper at www.nature.com/nature.

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Supplementary Information is linked to the online version of the paper at www.nature.com/nature.

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Author Contributions S.M.H. performed the single-molecule bead experiments; S.M.H. and J.J.L. performed the single-molecule fluorescence experiments; S.M.H. and M.T. performed the bulk-phase experiments; S.M.H., C.C.R. and A.M.v.O. designed the experiments, analysed the data and wrote the manuscript.

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METHODS

Single-molecule stretching and length measurements. Phage- λ DNA molecules were annealed and ligated to modified oligonucleotides to introduce a biotinylated fork on one end of the DNA and a digoxigenin moiety on the other end. The resulting DNA molecules were attached, by the 5' terminus of the bifurcated end, to the streptavidin-coated glass surface of a flow cell and, by the 3' end of the same strand, to a 2.8- μm -diameter anti-digoxigenin-coated paramagnetic bead (Dynal) (Supplementary Information). To prevent non-specific interactions between the beads and the surface, a 1-pN upward magnetic force was applied on the bead by positioning a permanent magnet above the flow cell. Beads were imaged with a charge-coupled-device camera at a time resolution of 500 ms and their positions were determined by particle-tracking software (Semasoph). Coordinated DNA synthesis in the flow cell was carried out by flowing gp4 helicase-primase (10 nM hexamers), T7 DNA polymerase (a purified 1:1 complex of gp5 and thioredoxin, 80 nM), and gp2.5 (750 nM) in buffer A (40 mM Tris-HCl (pH 7.5), 10 mM MgCl_2 , 10 mM DTT, 50 mM potassium glutamate (pH 7.5), 0.1 $\text{mg}\cdot\text{ml}^{-1}$ BSA), 600 μM dATP, 600 μM dCTP, 600 μM dGTP, 600 μM dTTP, 300 μM ATP and 300 μM CTP. Traces were corrected for instabilities in the flow by subtracting traces corresponding to tethers that were not enzymatically altered. Brownian motion and residual fluctuations resulted in a 300-bp error in DNA length measurement.

Fluorescence imaging. DNA was tethered at the forked end to functionalized coverslips as described above, but beads were omitted. Instead, the hydrodynamic drag on the DNA itself was used to extend the molecule. In the presence of 100 nM Sytox Orange dsDNA-specific stain (Invitrogen), the stretched DNA was imaged using through-objective TIRF microscopy (Olympus IX-71; $\times 60$, 1.45 numerical aperture). A continuous-wave 532-nm diode laser (CrystaLaser) was used to excite stain at power densities sufficiently low to minimize photo-induced cleavage of stained DNA over the timescale of an experiment. Protein and nucleotide concentrations were identical to those used to replicate the bead-tethered DNA molecules (see above). Single-molecule bead-tethering assays demonstrated that the kinetics of leading-strand synthesis and coordinated replication were not influenced by the presence of the stain (Supplementary Information).